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# **micca Documentation**

*Release 1.6.2*

**micca development team**

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micca (MICrobial Community Analysis) is a software pipeline for the processing of amplicon sequencing data, **from raw sequences to OTU tables, taxonomy classification and phylogenetic tree** inference. The pipeline can be applied to a range of highly conserved genes/spacers, such as **16S rRNA gene, Internal Transcribed Spacer (ITS)** and **28S rRNA**. micca is an open-source, GPLv3-licensed software.

- [Homepage](#)
- [Documentation \(latest\)](#)
- [Issues](#)
- [Github page](#)

Key features:

- supports **single-end** (Roche 454, Illumina MiSeq/HiSeq ,Ion Torrent) and **overlapping paired-end** reads (Illumina MiSeq/HiSeq);
- **multithread** de novo greedy, closed-reference, open-reference and swarm OTU picking protocols;
- **state-of-the-art taxonomic classification** algorithms (RDP and consensus-based classifier);
- fast and memory efficient **NAST** multiple sequence alignment (MSA);
- filters low quality sequences according to the maximum allowed **expected error (EE) rate %**;
- runs on **Linux, Mac OS X** and **MS Windows** (through Docker containers)
- **simple, easy to use.**

**Docker** images are available (compmetagen/micca) starting from version 1.2.2, see the documentation ( $\geq 1.3.0$ ) to learn how to use them. [Docker hub page](#).

**How to cite:** Davide Albanese, Paolo Fontana, Carlotta De Filippo, Duccio Cavalieri and Claudio Donati. **MICCA: a complete and accurate software for taxonomic profiling of metagenomic data**. Scientific Reports 5, Article number: 9743 (2015), doi:10.1038/srep09743, [Link](#). Dataset download: <ftp://ftp.fmach.it/metagenomics/micca/scirep/>.

micca wraps third party software packages and these **should be cited** if they are used:

- VSEARCH (doi: 10.7717/peerj.2584) used in `classify`, `filter`, `mergepairs`, `otu` and `msa` commands
- MUSCLE (doi: 10.1093/nar/gkh340) used in `msa` and `tree` commands
- FastTree (doi: 10.1371/journal.pone.0009490) used in the `tree` command
- Cutadapt (doi: 10.14806/ej.17.1.200) used in the `trim` command
- RDP classifier (doi: 10.1128/AEM.00062-07) used in the `classify` command
- swarm (doi: 10.7717/peerj.1420) used in the `otu` command



### Using Docker (that is, on MS Windows, Mac OS X and Linux!)

The easiest way to run micca is through [Docker](#). Docker works similarly to a virtual machine image, providing a container in which all the software has already been installed, configured and tested.

1. Install Docker for [Linux](#), [Mac OS X](#) or [Windows](#).
2. Run the Docker Quickstart Terminal (Mac OS X, Windows) or the docker daemon (Linux, `sudo service docker start`).
3. Download the latest version:

```
docker pull compmetagen/micca
```

4. Run an instance of the image, mounting the host working directory (e.g. `/Users/davide/micca`) on to the container working directory `/micca`:

```
docker run --rm -t -i -v /Users/davide/micca:/micca -w /micca compmetagen/micca /  
→bin/bash
```

You need to write something like `-v //c/Users/davide/micca:/micca` if you are in Windows or `-v /home/davide/micca:/micca` in Linux. The `--rm` option automatically removes the container when it exits.

5. Now you can use micca:

```
root@68f6784e1101:/micca# micca -h
```

---

**Note:** The RDP classifier is preinstalled in the Docker image, so you can check the software version by typing `echo $RDPPATH`

---

## Using pip

### On Ubuntu >= 12.04 and Debian >=7

We suggest to install the following packages through the package manager:

```
sudo apt-get update
sudo apt-get install build-essential python-numpy gcc gfortran python-dev libblas-dev
↳liblapack-dev cython pkg-config libfreetype6 libfreetype6-dev libpng-dev
```

Then, upgrade pip and install setuptools:

```
pip install --upgrade pip
pip install 'setuptools >=14.0'
```

Finally, install micca:

```
sudo pip install micca
```

### On Mac OS X

In Mac OS X, we recommend to install Python from [Homebrew](#):

1. Install [Xcode](#);
2. Install [Homebrew](#);
3. Make sure the environment variable `PATH` is properly setted in your `~/.bash_profile` or `~/.bashrc`:

```
.. code-block:: sh
```

```
export PATH=/usr/local/bin:$PATH
```

4. Install Python:

```
brew update
brew install python
```

Install the GNU Fortran and the NumPy package:

```
brew install gcc
pip install numpy
```

Finally, install micca:

```
sudo pip install micca
```

### Installation problems

- BIOM fatal error: 'numpy/arrayobject.h'. If the installation process returns a message like this:

```
biom/_filter.c:258:10: fatal error: 'numpy/arrayobject.h' file not found
#include "numpy/arrayobject.h"
      ^
```



```
1 error generated.
error: command 'clang' failed with exit status 1
```

then you need to run:

```
pip install --global-option=build_ext --global-option="-I/usr/local/lib/python2.7/
↳site-packages/numpy/core/include/" biom-format
```

After that you can install the micca package.

## Install micca from source

In order to install micca from sources (with the standard procedure `python setup.py install`), in addition to Python ( $\geq 2.7$ ) and NumPy ( $\geq 1.8.0$ ), the following Python packages must be installed:

- SciPy  $\geq 0.13.0$
- Pandas  $\geq 0.17.0$
- matplotlib  $\geq 1.3.0$
- Biopython  $\geq 1.50$
- cutadapt  $\geq 1.9$
- biom-format  $\geq 1.3.1$

The easiest way to install these packages is to is using pip:

```
sudo pip install 'scipy >=0.13.0' 'pandas >=0.17.0' 'matplotlib >=1.3.0' 'biopython >
↳= 1.50' 'cutadapt >=1.9' 'biom-format >=1.3.1'
```

Download the latest version from <https://github.com/compmetagen/micca/releases> and complete the installation:

```
tar -zxvf micca-X.Y.Z.tar.gz
sudo python setup.py install
```

## If you don't have root access

Install micca in a local directory by specifying the `--prefix` argument. Then you need to set the environment variable `PYTHONPATH`:

```
python setup.py install --prefix=/path/to/modules
export PYTHONPATH=$PYTHONPATH:/path/to/modules/lib/python{version}/site-packages
```

**Note:** In order to export the variable permanently add the command at the bottom of your `~/.bash_profile` or `~/.bashrc` file.

## Testing the installation

```
micca -h
```

## Install RDP classifier (optional)

The RDP Classifier is a naive bayesian classifier for taxonomic assignments (<http://sourceforge.net/projects/rdp-classifier/>). The RDP classifier can be used in the *classify* command (option `-m/--method rdp`).

**Warning:** Only RDP Classifier version >2.8 is supported. Install the standard Java or Java compatible runtime (sudo apt-get install default-jre in Ubuntu/Debian or go to the Oracle Java homepage for OS X)

Download and unzip the file (RDP classifier 2.11 2015-09-14):

```
wget https://sourceforge.net/projects/rdp-classifier/files/rdp-classifier/rdp_
↪classifier_2.11.zip
unzip rdp_classifier_2.11.zip
```

Now you must set the environment variable RDPPATH by typing:

```
$ export RDPPATH=/path-to-rdp-classifier/rdp_classifier_2.11/
```

e.g. export RDPPATH=/Users/David/rdp\_classifier\_2.11.

---

**Note:** In order to export the variable permanently add the latest command at the bottom of your `.bashrc` file.

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## CHAPTER 2

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### Run micca in 6 steps

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Open a terminal, download the sample data and prepare the working directory:

```
wget ftp://ftp.fmach.it/metagenomics/micca/examples/mwanihana.tar.gz
tar -zxvf mwanihana.tar.gz
cd mwanihana
```

Now we can run micca:

```
# merge the samples
micca merge -i fastq/*.fastq -o merged.fastq
# trim primers
micca trim -i merged.fastq -o trimmed.fastq -w AGAGTTTGATCMTGGCTCAG -r_
↪GTGCCAGCAGCCGCGGTAA -W
# filter low quality reads and truncate at 350 bp
micca filter -i trimmed.fastq -o filtered.fasta -e 0.5 -m 350 -t
# denovo OTU picking protocol
micca otu -i filtered.fasta -o denovo_greedy_otus -d 0.97 -c -t 4
# classify taxonomies using RDP classifier
micca classify -m rdp -i denovo_greedy_otus/otus.fasta -o denovo_greedy_otus/taxa.txt
# export the BIOM file
micca tobiom -i denovo_greedy_otus/otutable.txt -o denovo_greedy_otus/tables.biom -t_
↪denovo_greedy_otus/taxa.txt
```



---

## Supported databases

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### 16S rRNA

- Greengenes [ftp://greengenes.microbio.me/greengenes\\_release/](ftp://greengenes.microbio.me/greengenes_release/)
- Greengenes *Core Set* (useful for NAST) [http://greengenes.lbl.gov/Download/Sequence\\_Data/Fasta\\_data\\_files/core\\_set\\_aligned.fasta.imputed](http://greengenes.lbl.gov/Download/Sequence_Data/Fasta_data_files/core_set_aligned.fasta.imputed)
- QIIME-formatted SILVA <https://www.arb-silva.de/download/archive/qiime/>

### ITS

- UNITE <https://unite.ut.ee/repository.php> (QIIME releases)

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**Note:** In the QIIME releases of the UNITE database the `_s` in the filename (e.g. `sh_qiime_release_s_DD.MM.YYY.zip`) specifies that the database includes singletons.

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## Single-end sequencing

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This tutorial describes a standard micca pipeline for the analysis of single-end amplicon data. This pipeline is intended for different platforms, such as **Roche 454**, **Illumina MiSeq/HiSeq** and **Ion Torrent**. Although this tutorial explains how to apply the pipeline to **16S rRNA** amplicons, it can be adapted to others markers gene/spacers, e.g. **Internal Transcribed Spacer (ITS)** or **28S**.

### Table of Contents

- *Dataset download*
- *Merge files*
- *Primer trimming*
- *Quality filtering*
- *OTU picking*
- *Assign taxonomy*
- *Infer the phylogenetic tree*
- *Build the BIOM file*
- *Import and analyze the data into R and phyloseq*
- *Compute basic OTU table statistics, rarefy and summarize OTU tables by taxa using micca*

## Dataset download

The dataset used in this tutorial is taken from the Barelli et al. paper *Habitat fragmentation is associated to gut microbiota diversity of an endangered primate: implications for conservation* (<https://doi.org/doi:10.1038/srep14862>). The dataset contains only a subset of the entire study (Mwanihana samples only) for a total of **15 samples** (in FASTQ format) and **235179 16S rRNA amplicon reads** (V1-V3 hypervariable regions, 27-Forward

5'-AGAGTTTGATCMTGGCTCAG, 533-Reverse 5'-TTACCGCGGCTGCTGGCAC). The 454 pyrosequencing was carried out on the GS FLX+ system using the XL+ chemistry.

Open a terminal, download the data and prepare the working directory:

```
wget ftp://ftp.fmach.it/metagenomics/micca/examples/mwanihana.tar.gz
tar -zxvf mwanihana.tar.gz
cd mwanihana
```

## Merge files

Now the FASTQ files must be merged in a single file. This operation can be performed with the *merge* command. Sample names will be included into the sequence identifiers.

```
micca merge -i fastq/*.fastq -o merged.fastq
```

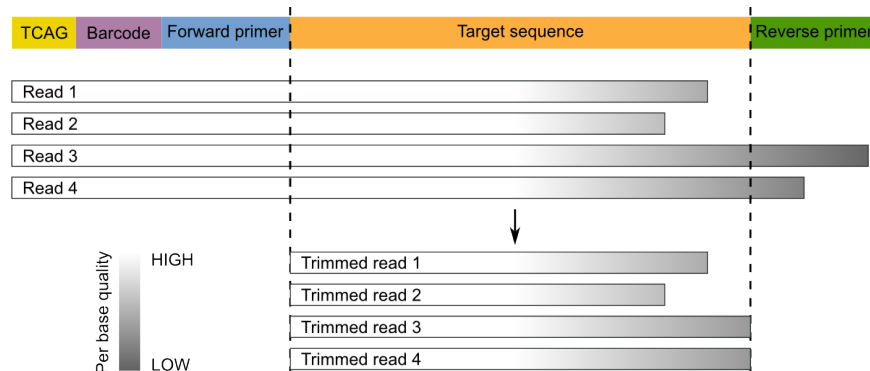
**Note:** The *merge* command works with FASTQ or FASTA files. If your sequences are in a different format (e.g. SFF or FASTA+QUAL) use *convert* to convert them.

**Warning:** In the case of multiplexed reads (with 5' barcode sequences) use *split* instead of *merge*. This command will perform demultiplexing and merging at the same time.

**Warning:** In the case of overlapping paired-end reads go to *Paired-end sequencing*.

## Primer trimming

Segments which match PCR primers should be now removed. Typical Roche 454 reads start with a sequence key (e.g. TCAG) followed by the barcode (if it was not previously removed) and the forward primer. For these types of data (and in general, for single-end sequencing) we recommend to **trim both forward reverse primers and discard reads that do not contain the forward primer**. Moreover, sequence preceding (for the forward) or succeeding (for the reverse, if found) primers should be removed:



These operations can be performed with the *trim* command:



```
micca trim -i merged.fastq -o trimmed.fastq -w AGAGTTGATCMTGGCTCAG -r_
↪GTGCCAGCAGCCGCGGTAA -W
```

The option `-W/--duforward` ensures that reads that do not contain the forward primer will be discarded.

**Warning:** Do not use the `-R/--dureverse` with single-end reads.

**Note:** The `trim` command supports IUPAC nucleotide codes and multiple primers. With the option `-c/--searchrc` the command searches reverse complement primers too. `trim` works with FASTQ or FASTA files.

## Quality filtering

Producing high-quality OTUs requires high-quality reads. The `filter` command filters sequences according to the maximum allowed expected error (EE) rate % (see [Quality filtering strategy in micca](#)). We recommend values  $\leq 1\%$ . Moreover, to obtain good results in clustering (see `otu`), reads should be **truncated at the same length** when they cover partial amplicons or if quality deteriorates towards the end (common when you have long amplicons in single-end sequencing).

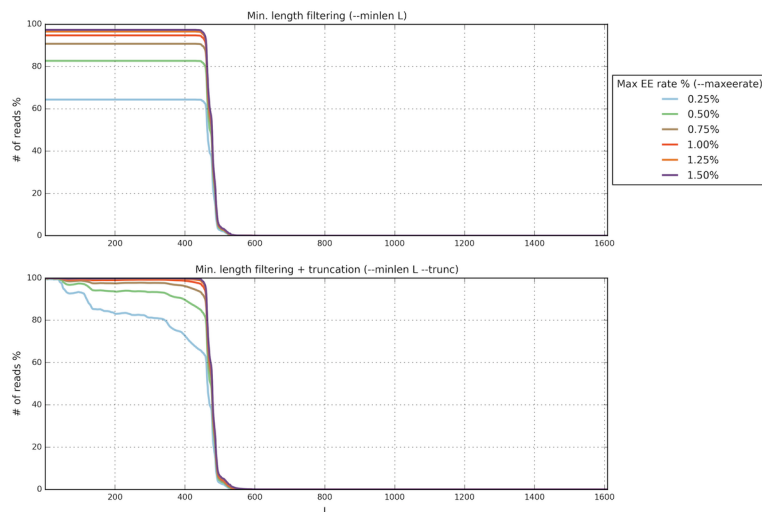
**Warning:** Parameters for the `filter` command should be chosen using the command `filterstats`.

## Choosing parameters for filtering

The command `filterstats` reports the fraction of reads that would pass for each specified maximum expected error (EE) rate %:

```
micca filterstats -i trimmed.fastq -o filterstats
```

Open the PNG file `filterstats/stats_plot.png`:



In this case we are interested in the plot below (minimum length filtering + truncation). A truncation length of **350** and a maximum error rate of **0.5%** seems to be a good compromise between read length, expected error rate and number of reads remaining. Inspecting the file `filterstats/truncrlen_stats.txt`, you can see that more than **92%** reads will pass the filter:

L	0.25	0.5	0.75	1.0	1.25	1.5
...						
349	78.905	92.472	97.425	99.135	99.705	99.897
350	78.639	92.385	97.389	99.126	99.704	99.896
351	78.369	92.300	97.357	99.116	99.700	99.892
...						

---

**Note:** To obtain general sequencing statistics, run `stats`.

---

## Filter sequences

Now we can run the `filter` command with the selected parameters:

```
micca filter -i trimmed.fastq -o filtered.fasta -e 0.5 -m 350 -t
```

---

**Note:** The maximum number of allowed Ns after truncation can be also specified in `filterstats` and in `filter`.

---

## OTU picking

To characterize the taxonomic structure of the samples, the sequences are now organized into **Operational Taxonomic Units (OTUs)** at varying levels of identity. An identity of **97%** represent the common working definition of bacterial species. The `otu` command implements several state-of-the-art approaches for OTU clustering, but in this tutorial we will focus on the **de novo greedy clustering** (see *OTU picking in micca*):

```
micca otu -i filtered.fasta -o denovo_greedy_otus -d 0.97 -c -t 4
```

The `otu` command returns several files in the output directory, including the **OTU table** (`otutable.txt`) and a FASTA file containing the **representative sequences** (`otus.fasta`).

---

**Note:** See *OTU picking in micca* to see how to apply the **swarm de novo**, **closed-reference** and the **open-reference** OTU picking strategies to these data.

---

## Assign taxonomy

Now we can assign taxonomy to each representative sequence using the `classify` command. In this tutorial we use the RDP (<https://doi.org/10.1128/AEM.00062-07>) classifier.

```
micca classify -m rdp -i denovo_greedy_otus/otus.fasta -o denovo_greedy_otus/taxa.txt
```

`classify` returns a taxonomy file like this:

```
DENOVO1 Bacteria;Firmicutes;Clostridia;Clostridiales
DENOVO2 Bacteria;Bacteroidetes;Bacteroidia;Bacteroidales
DENOVO3 Bacteria;Verrucomicrobia;Opitutae
DENOVO4 Bacteria;Bacteroidetes;Bacteroidia;Bacteroidales
...
```

## Infer the phylogenetic tree

These steps are necessary if you want to use phylogenetic-based metrics such as the UniFrac distance (<https://doi.org/10.1128/AEM.01996-06>) in the downstream analysis.

### Multiple Sequence Alignment (MSA)

The *msa* command provides two approaches for MSA: MUSCLE (<https://doi.org/10.1093/nar/gkh340>) (de novo alignment) and Nearest Alignment Space Termination (NAST) (<https://doi.org/10.1093/nar/gkl244>) (which uses a template alignment). In this tutorial we will use the NAST alignment method. For 16S rRNA sequences, a good template alignment is the Greengenes Core Set:

```
wget ftp://ftp.fmach.it/metagenomics/micca/dbs/core_set.tar.gz
tar -zxvf core_set.tar.gz
```

At this point we can run the *msa* command:

```
micca msa -m nast -i denovo_greedy_otus/otus.fasta -o denovo_greedy_otus/msa.fasta --
↳nast-template core_set_aligned.fasta.imputed --nast-threads 4
```

### Build the phylogenetic tree

At this point we can build the phylogenetic tree from the MSA using *tree*:

```
micca tree -i denovo_greedy_otus/msa.fasta -o denovo_greedy_otus/tree.tree
```

---

**Note:** The output tree is in [Newick format](#).

---

### Midpoint rooting

UniFrac metrics require phylogenetic trees to be rooted. The tree can be rooted (in this case at midpoint between the two most distant tips of the tree) using the *root* command:

```
micca root -i denovo_greedy_otus/tree.tree -o denovo_greedy_otus/tree_rooted.tree
```

---

**Note:** Tree can also be rooted with the outgroup clade containing selected targets, see [root](#).

---

## Build the BIOM file

The [Biological Observation Matrix \(BIOM\)](#) is a common format for representing OTU tables and metadata and is the core data type for downstream analyses in [QIIME](#) and in [phyloseq](#). `tobiom` converts the OTU table and the taxonomy table produced by the previous steps to the BIOM format. In addition, the *Sample data* can be added:

```
micca tobiom -i denovo_greedy_otus/otutable.txt -o denovo_greedy_otus/tables.biom -t ↵
↵denovo_greedy_otus/taxa.txt -s sampledata.txt
```

## Import and analyze the data into R and phyloseq

See *Import and analyze the data into R and phyloseq*

## Compute basic OTU table statistics, rarefy and summarize OTU tables by taxa using micca

See *Compute basic OTU table statistics, rarefy and summarize OTU tables by taxa using micca*

---

## Paired-end sequencing

---

This tutorial describes a standard micca pipeline for the analysis of overlapping paired-end Illumina data. This pipeline is intended for different platforms, such as **Illumina MiSeq** and **Illumina HiSeq**. Although this tutorial explains how to apply the pipeline to **16S rRNA** amplicons, it can be adapted to others markers gene/spacers, e.g. **Internal Transcribed Spacer (ITS)** or **28S**.

### Table of Contents

- *Dataset download*
- *Merge paired-end sequences*
- *Primer trimming*
- *Quality filtering*
- *OTU picking, taxonomy assignment, phylogenetic tree inference and downstream analysis*

## Dataset download

This paired-end 16S rRNA dataset contains **3 samples** in FASTQ format (V3-V4 region, 341-Forward 5'-CCTACGGGNGGCWGCAG-3', 805-Reverse 5'-GACTACNVGGGTWTCTAATCC-3'). The 300-bp paired-end sequencing was carried out on an Illumina MiSeq.

Open a terminal, download the data and prepare the working directory:

```
wget ftp://ftp.fmach.it/metagenomics/micca/examples/pairedend.tar.gz
tar -zxvf pairedend.tar.gz
cd pairedend
```

## Merge paired-end sequences

Now the paired sequences must be merged to obtain consensus sequences (sometimes called *assembly*). This operation can be performed with the `mergepairs` command. After the merging of the paired reads, the `mergepairs` command merges the different samples in a single file where sample names are appended to the sequence identifier, as in `merge` and `split`. Passing the forward files only reverse file names will be constructed by replacing the string `_R1` in the forward file name with `_R2` (typical in Illumina file names, see options `-p/--pattern` and `-e/--repl`).

Since the sequenced region is about of 464-bp (805-341) and the reads are of 300-bp, the overlap region is quite large (~136 bp), as rule of thumb we set a minimum overlap length of 100 and maximum number of allowed mismatches of about 1/3, say 32:

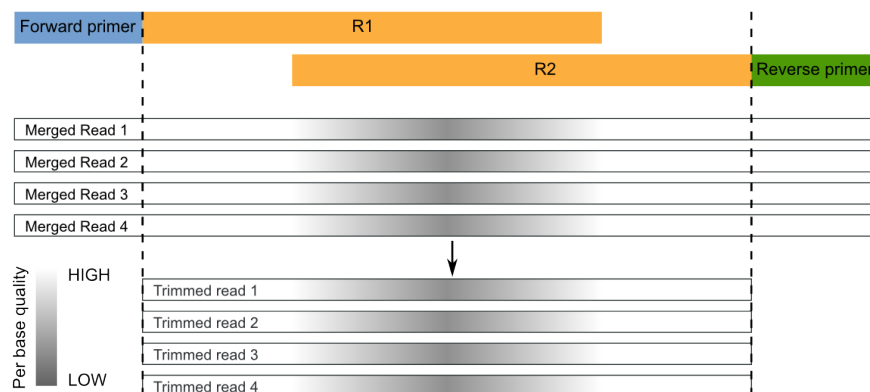
```
micca mergepairs -i fastq/*_R1*.fastq -o merged.fastq -l 100 -d 32
```

**Note:** Starting from micca 1.6.0 staggered read pairs (staggered pairs are pairs where the 3' end of the reverse read has an overhang to the left of the 5' end of the forward read) will be merged by default. To override this feature (and therefore to discard staggered alignments) set the `-n/--nostagger` option.

**Note:** `mergepairs` works with FASTQ files only.

## Primer trimming

Segments which match PCR primers should be now removed. For Illumina paired-end (already merged) reads, we recommend to **trim both forward and reverse primers and discard reads that do not contain the forward OR the reverse primer**. Moreover, sequence preceding (for the forward) or succeeding (for the reverse) primers should be removed:



These operations can be performed with the `trim` command:

```
micca trim -i merged.fastq -o trimmed.fastq -w CCTACGGGNGGCWGCAG -r_
↳GACTACNVGGGTWCTAATCC -W -R -c
```

The option `-W/--duforward` and `-R/--dureverse` ensures that reads that do not contain the forward or the reverse primer will be discarded. With the option `-c/--searchrc` the command searches reverse complement primers too.

## Quality filtering

Producing high-quality OTUs requires high-quality reads. *filter* filters sequences according to the maximum allowed expected error (EE) rate % (see *Quality filtering strategy in micca*). We recommend values  $\leq 1\%$ .

For paired-end reads, we recommend to merge pairs first, then quality filter using a maximum EE threshold with **no length truncation**.

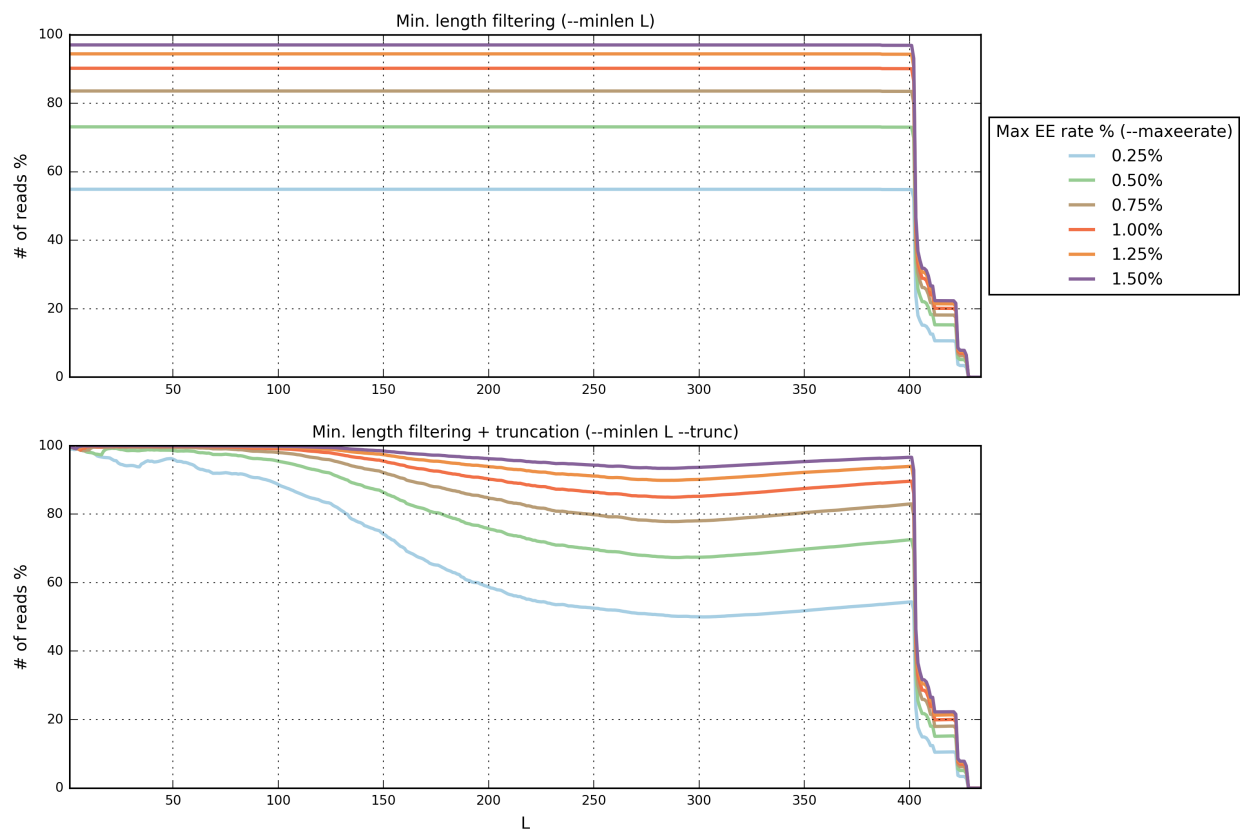
**Warning:** Parameters for the *filter* command should be chosen using the tool *filterstats*.

## Choosing parameters for filtering

The command *filterstats* reports the fraction of reads that would pass for each specified maximum expected error (EE) rate %:

```
micca filterstats -i trimmed.fastq -o filterstats
```

Open the PNG file `filterstats/stats_plot.png`:



In this case we are interested in the plot on top (minimum length filtering only). A truncation length of **400** and a maximum error rate of **0.5%** seems to be a good compromise between the expected error rate and the number of reads remaining. Inspecting the file `filterstats/minlen_stats.txt`, you can see that more than **73%** reads will pass the filter:

```
L          0.25    0.5    0.75    1.0    1.25    1.5
...
399  54.801  73.016  83.476  90.107  94.312  96.917
400  54.799  73.013  83.473  90.104  94.309  96.914
401  54.781  72.993  83.452  90.080  94.285  96.890
...
```

---

**Note:** To obtain general sequencing statistics, run *stats*.

---

## Filter sequences

Now we can run the *filter* command with the selected parameters:

```
micca filter -i trimmed.fastq -o filtered.fasta -e 0.5 -m 400
```

---

**Note:** The maximum number of allowed Ns after truncation can be also specified in *filterstats* and in *filter*.

---

## OTU picking, taxonomy assignment, phylogenetic tree inference and downstream analysis

See *Single-end sequencing* starting from *OTU picking*.



---

## Import and analyze the data into R and phyloseq

---



---

**Note:** This tutorial requires *Single-end sequencing* to be done.

---



---

**Note:** This tutorial requires R and phyloseq (tested on R v3.2.1 and phyloseq v1.14.0) to be installed in your system.

---

We can import the micca processed data (the BIOM file, the phylogenetic tree and the representative sequences) into the R environment using the `phyloseq` library. From the phyloseq homepage:

“The phyloseq package is a tool to import, store, analyze, and graphically display complex phylogenetic sequencing data that has already been clustered into Operational Taxonomic Units (OTUs), especially when there is associated sample data, phylogenetic tree, and/or taxonomic assignment of the OTUs.”

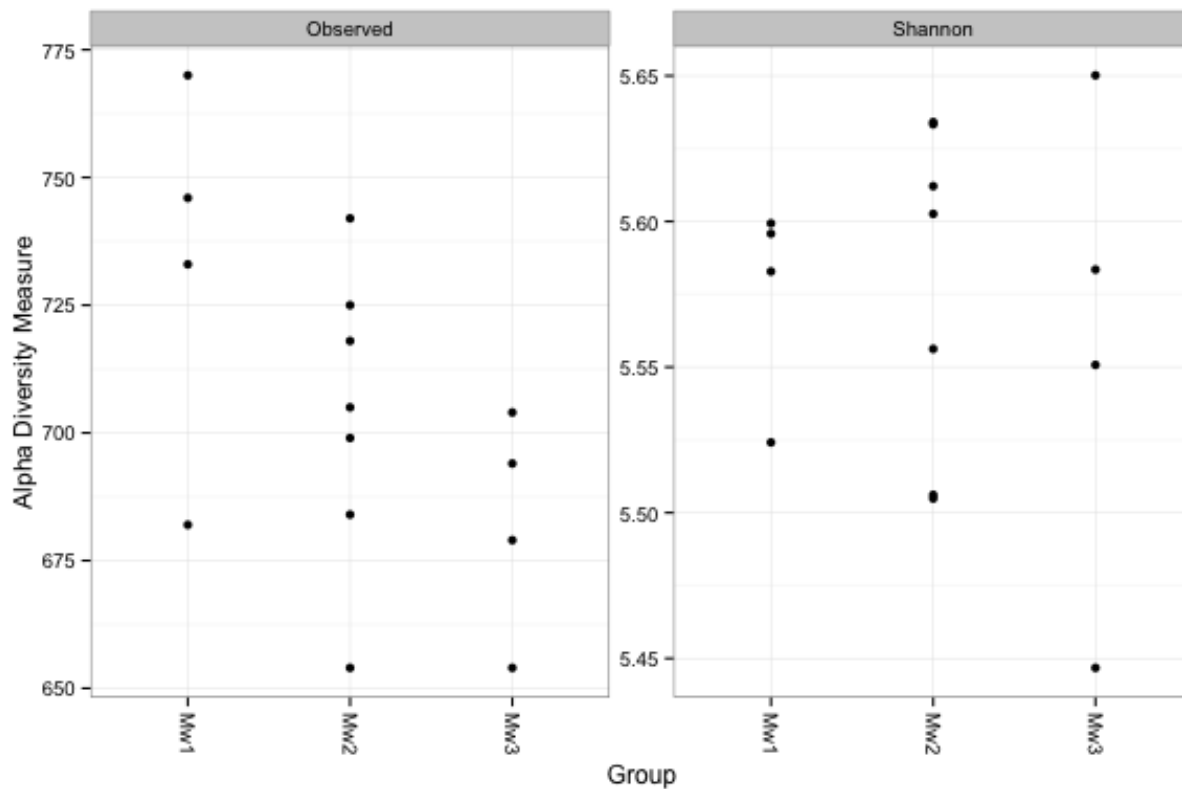
The `import_biom()` function allows to simultaneously import the BIOM file and an associated phylogenetic tree file and reference sequence file.

```
> library("phyloseq")
> library("ggplot2")
> theme_set(theme_bw())
> setwd("denovo_greedy_otus") # set the working directory
> ps = import_biom("tables.biom", treefilename="tree_rooted.tree",
+ refseqfilename="otus.fasta")
> ps
phyloseq-class experiment-level object
otu_table() OTU Table: [ 1332 taxa and 15 samples ]
sample_data() Sample Data: [ 15 samples by 2 sample variables ]
tax_table() Taxonomy Table: [ 1332 taxa by 6 taxonomic ranks ]
phy_tree() Phylogenetic Tree: [ 1332 tips and 1331 internal nodes ]
refseq() DNASTringSet: [ 1332 reference sequences ]
```

At this point we can compute the number of OTUs and the Shannon diversity index after the rarefaction:

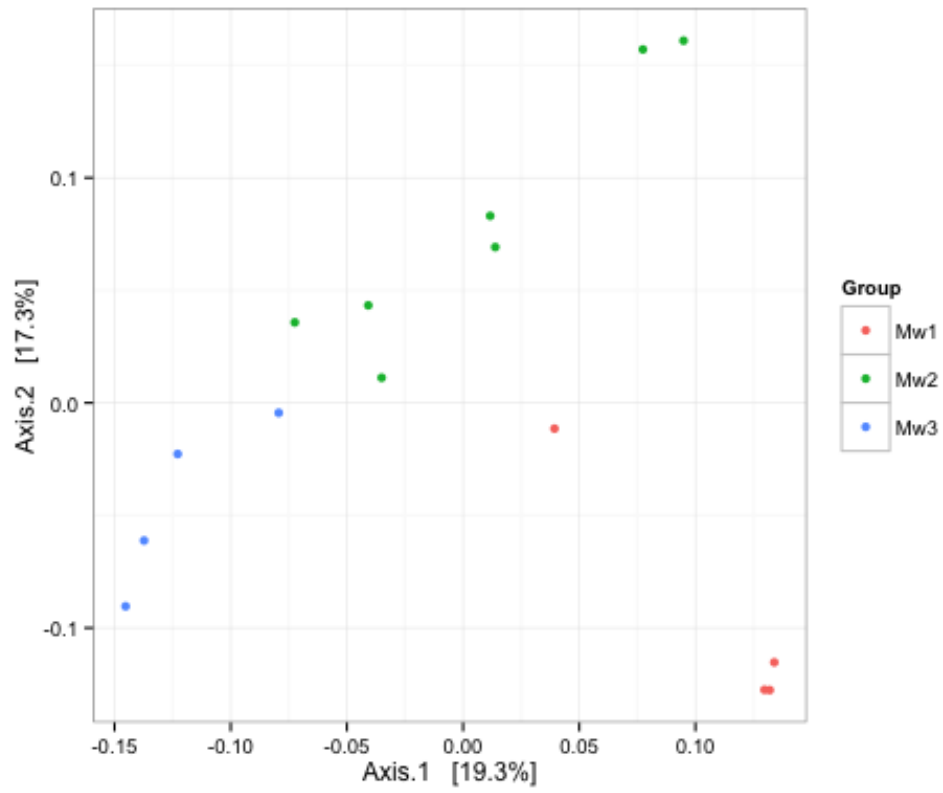
```
> # rarefy without replacement
> ps.rarefied = rarefy_even_depth(ps, rngseed=1, sample.size=0.9*min(sample_sums(ps)),
↪ replace=F)
```

```
> # plot alpha diversity indexes  
> plot_richness(ps.rarefied, x="Group", measures=c("Observed", "Chao1", "Shannon"))
```



Finally, we can plot the PCoA on the unweighted UniFrac distance of samples:

```
> # PCoA plot on the unweighted UniFrac distance  
> ordination = ordinate(ps.rarefied, method="PCoA", distance="unifrac", weighted=F)  
> plot_ordination(ps.rarefied, ordination, color="Group") + theme(aspect.ratio=1)
```





---

## Compute basic OTU table statistics, rarefy and summarize OTU tables by taxa using micca

---

**Note:** This tutorial requires *Single-end sequencing* to be done.

---

The command `tablestats` reports a sample summary, an OTU summary and the rarefaction curves for the input OTU table:

```
micca tablestats -i otutable.txt -o tablestats
```

Inspecting the file `tablestats/tablestats_samplesumm.txt` you can see that the less abundant sample contains 9053 reads:

```
Sample Depth NOTU NSingle
Mw_03  9053  716  142
Mw_02  9947  760  166
Mw_12 10843  792  168
...    ...    ...    ...
```

**Note:** Rarefaction curves can be inspected through `tablestats/tablestats_rarecurve.txt` and `tablestats/tablestats_rarecurve_plot.png`.

---

To compare different samples, the OTU table must be subsampled (*rarefied*) using the command `tablerare`. In this case we are interested in rarefy the table with the depth of the less abundant sample (Mw\_03):

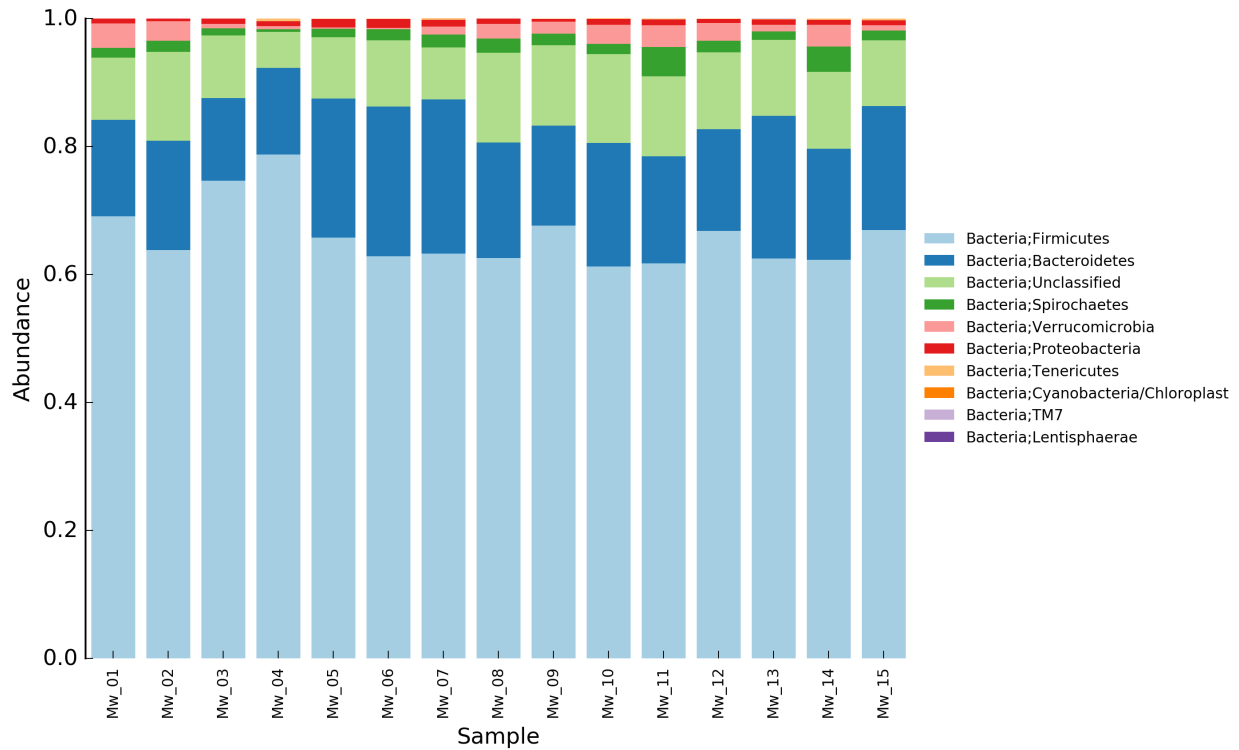
```
micca tablerare -i otutable.txt -o otutable_rare.txt -d 9053
```

Now we can summarize communities by their taxonomic composition. The `tabletotax` creates in the output directory a table for each taxonomic level (`taxtable1.txt`, ..., `taxtableN.txt`). OTU counts are summed together if they have the same taxonomy at the considered level.

```
micca tabletotax -i otutable_rare.txt -t taxa.txt -o taxtables
```

Finally, we can generate a relative abundance bar plot from generated taxa tables, using the command *tablebar*. In this case only the bar plot relative to the taxonomy level 2 (phylum) will be generated:

```
micca tablebar -i taxtables/taxtable2.txt -o taxtables/taxtable2.png
```



---

## Picking OTUs for use in PICRUSt

---

**PICRUSt** (doi: 10.1038/nbt.2676) is a software designed to predict metagenome functional content from marker gene (e.g., 16S rRNA) surveys and full genomes. This tutorial covers how to pick OTUs from 16S rRNA sequences data to use with PICRUSt.

**Note:** Requires *Quality filtering in Single-end sequencing* to be done and the PICRUSt software to be installed in your system. Warning: PICRUSt 1.0.0 requires the biom-format package v1.3.1 to be installed in your system (from the command line run: `pip install biom-format==1.3.1`, for more information see <http://biom-format.org/>).

PICRUSt requires an *Closed-reference* OTU table computed against the Greengenes reference (clustered at 97% identity). Download the reference database (Greengenes, version 2013/05), clustered at 97% identity:

```
wget ftp://ftp.fmach.it/metagenomics/micca/dbs/gg_2013_05.tar.gz
tar -zxvf gg_2013_05.tar.gz
```

Run the micca closed-reference protocol:

```
micca otu -m closed_ref -i filtered.fasta -o closed_ref_otus -r 97_otus.fasta -d 0.97
↪ -t 4
cd closed_ref_otus
```

Report the sample summary:

```
micca tablestats -i otutable.txt -o tablestats
head tablestats/tablestats_samplesumm.txt
```

Sample	Depth	NOTU	NSingle
Mw_03	1084	132	39
Mw_06	1387	122	27
Mw_11	1485	155	44
Mw_07	1528	150	36
Mw_01	1537	143	35
Mw_15	1565	144	35
Mw_14	1610	149	42

Mw_02	1670	143	43
Mw_12	1710	153	54

Rarefy the OTU table for the PICRUSt analysis is always a good idea (see <https://groups.google.com/forum/#!topic/picrust-users/ev5uZGUIPrQ>), so we will rarefy the table at 1084 sequences per sample using *tablerare*:

```
micca tablerare -i otutable.txt -o otutable_rare.txt -d 1084
```

Convert the rarefied OTU table into the BIOM format replacing the OTU IDs with the original sequence IDs using the *tobiom* command:

```
micca tobiom -i otutable_rare.txt -o tables.biom -u otuids.txt
```

Normalize the OTU table by dividing each OTU by the known/predicted 16S copy number abundance:

```
normalize_by_copy_number.py -i tables.biom -o normalized_otus.biom
```

Create the final metagenome functional predictions:

```
predict_metagenomes.py -i normalized_otus.biom -o metagenome_predictions.biom
```

Now you can analyze the PICRUSt predicted metagenome: [http://picrust.github.io/picrust/tutorials/downstream\\_analysis.html#downstream-analysis-guide](http://picrust.github.io/picrust/tutorials/downstream_analysis.html#downstream-analysis-guide).



## CHAPTER 9

---

### Quality filtering strategy in micca

---

TODO



## OTU picking in micca

To characterize the taxonomic structure of the samples, the sequences are organized into [Operational Taxonomic Units \(OTUs\)](#) at varying levels of identity. An identity of **97%** represent the common working definition of bacterial species. The `otu` command assigns similar sequences (marker genes such as 16S rRNA and the fungal ITS region) to operational taxonomic units (OTUs). The command `otu` wraps [VSEARCH](#) for low-level clustering, chimera detection and searching operations.

The `otu` command returns in a single directory 5 files:

**otutable.txt** TAB-delimited file, containing the number of times an OTU is found in each sample (OTU x sample, see [Supported file formats](#)):

```
OTU      Mw_01 Mw_02 Mw_03 ...
DENOVO1 151   178   177   ...
DENOVO2 339   181   142   ...
DENOVO3 533   305   63    ...
...      ...   ...   ...   ...
```

**otus.fasta** FASTA containing the representative sequences (OTUs):

```
>DENOVO1
GACGAACGCTGGCGGCGTGCCTAACACATGCAAGTCGAACGGGG...
>DENOVO2
GATGAACGCTAGCTACAGGCTTAACACATGCAAGTCGAGGGGCA...
>DENOVO3
AGTGAACGCTGGCGACGTGGTTAAGACATGCAAGTCGAGCGGTA...
...
```

**otuids.txt** TAB-delimited file which maps the OTU ids to original sequence ids:

```
DENOVO1 IS0AYJS04JQKIS;sample=Mw_01
DENOVO2 IS0AYJS04JL6RS;sample=Mw_01
DENOVO3 IS0AYJS04H4XNN;sample=Mw_01
...
```

**hits.txt** TAB-separated file, three-columns, where each column contains: the matching sequence, the representative (seed) and the identity (if available), see *Definition of identity*:

```
IS0AYJS04JE658;sample=Mw_01; IS0AYJS04I4XYN;sample=Mw_01 99.4
IS0AYJS04JPH34;sample=Mw_01; IS0AYJS04JVUBC;sample=Mw_01 98.0
IS0AYJS04I67XN;sample=Mw_01; IS0AYJS04JVUBC;sample=Mw_01 99.7
...
```

**otuschim.fasta** (only for 'denovo\_greedy', 'denovo\_swarm' and 'open\_ref' methods, when `-c/` `--rmchim` is specified) FASTA file containing the chimeric otus.

**Warning:** Trimming the sequences to a fixed position before clustering is strongly recommended when they cover partial amplicons or if quality deteriorates towards the end (common when you have long amplicons and single-end sequencing), see *Quality filtering*.

---

**Note:** De novo OTUs are renamed to `DENOVO [N]` and reference OTUs to `REF [N]`.

---

## Clustering strategies

*otu* implements several state-of-the-art clustering strategies:

- *De novo greedy*
- *Closed-reference*
- *Open-reference*
- *De novo swarm*

### De novo greedy

In denovo greedy clustering (parameter `--method denovo_greedy`), sequences are clustered without relying on an external reference database, using an approach similar to the UPARSE pipeline (<https://doi.org/10.1038/nmeth.2604>) and tested in <https://doi.org/10.7287/peerj.preprints.1466v1>. *otu* includes in a single command dereplication, clustering and chimera filtering:

1. Dereplication. Predict sequence abundances of each sequence by dereplication, order by abundance and discard sequences with abundance value smaller than `DEREP_MINSIZE` (option `--derep-minsize` recommended value 2);
2. Greedy clustering. Distance (DGC) and abundance-based (AGC) strategies are supported (option `--greedy`, see <https://doi.org/10.1186/s40168-015-0081-x> and <https://doi.org/10.7287/peerj.preprints.1466v1> ). Therefore, the candidate representative sequences are obtained;
3. Chimera filtering (optional). Remove chimeric sequences from the representatives performing a de novo chimera detection (option `--rmchim`, recommended);
4. Map sequences. Map sequences to the representatives.

Example (requires *Quality filtering* in *Single-end sequencing* to be done):

```
micca otu -m denovo_greedy -i filtered.fasta -o denovo_greedy_otus -d 0.97 -c -t 4
```

## Closed-reference

Sequences are clustered against an external reference database and reads that could not be matched are discarded. Example (requires *Quality filtering* in *Single-end sequencing* to be done):

Download the reference database (Greengenes), clustered at 97% identity:

```
wget ftp://ftp.fmach.it/metagenomics/micca/dbs/gg_2013_05.tar.gz
tar -zxvf gg_2013_05.tar.gz
```

Run the closed-reference protocol:

```
micca otu -m closed_ref -i filtered.fasta -o closed_ref_otus -r 97_otus.fasta -d 0.97 -t 4
```

Simply perform a sequence ID matching with the reference taxonomy file (see *classify*):

```
cd closed_ref_otus
micca classify -m otuid -i otuids.txt -o taxa.txt -x ../97_otu_taxonomy.txt
```

## Open-reference

Open-reference clustering (*open\_ref*): sequences are clustered against an external reference database (as in *Closed-reference*) and reads that could not be matched are clustered with the *De novo greedy* protocol. Example (requires *Quality filtering* in *Single-end sequencing* to be done):

Download the reference database (Greengenes), clustered at 97% identity:

```
wget ftp://ftp.fmach.it/metagenomics/micca/dbs/gg_2013_05.tar.gz
tar -zxvf gg_2013_05.tar.gz
```

Run the open-reference protocol:

```
micca otu -m open_ref -i filtered.fasta -o open_ref_otus -r 97_otus.fasta -d 0.97 -t 7 -c
```

Run the VSEARCH-based consensus classifier or the RDP classifier (see *classify*):

```
cd open_ref_otus
micca classify -m cons -i otus.fasta -o taxa.txt -r ../97_otus.fasta -x ../97_otu_taxonomy.txt -t 4
```

## De novo swarm

In denovo swarm clustering (doi: 10.7717/peerj.593, doi: 10.7717/peerj.1420, <https://github.com/torognes/swarm>, parameter `--method denovo_swarm`), sequences are clustered without relying on an external reference database. From <https://github.com/torognes/swarm>:

The purpose of swarm is to provide a novel clustering algorithm that handles massive sets of amplicons. Results of traditional clustering algorithms are strongly input-order dependent, and rely on an arbitrary global clustering threshold. swarm results are resilient to input-order changes and rely on a small local

linking threshold  $d$ , representing the maximum number of differences between two amplicons. `swarm` forms stable, high-resolution clusters, with a high yield of biological information.

`otu` includes in a single command dereplication, clustering and de novo chimera filtering:

1. Dereplication. Predict sequence abundances of each sequence by dereplication, order by abundance and discard sequences with abundance value smaller than `DEREP_MINSIZE` (option `--derep-minsize` recommended value is 1, i.e. no filtering);
2. Swarm clustering. Number of differences 1 and the fastidious option are recommended (`--swarm-differences 1 --swarm-fastidious`).
3. Chimera filtering (optional). Remove chimeric sequences from the representatives performing a de novo chimera detection (option `--rmchim`);

**Warning:** Removing ambiguous nucleotides (N) (with the option `--maxns 0` in *filter*) is mandatory if you use the de novo swarm clustering method.

Example (requires *Primer trimming* in *Single-end sequencing* to be done):

```
micca filter -i trimmed.fastq -o filtered.fasta -e 0.5 -m 350 -t --maxns 0
micca otu -m denovo_swarm -i filtered.fasta -o otus_denovo_swarm -c --minsize 1 --
->swarm-fastidious -t 4
```

## Definition of identity

In `micca`, the pairwise identity (except for de novo swarm) is defined as the edit distance excluding terminal gaps (same as in USEARCH and BLAST):

$$\frac{\text{\# matching columns}}{\text{alignment length} - \text{terminal gaps}}$$

---

## Supported file formats

---

### Sequence files

**FASTA** and **FASTQ** Sanger/Illumina 1.8+ format (phred+33) formats are supported. `micca` provides the `convert` command to convert between sequence file formats.

### Taxonomy files

Taxonomy files map sequence IDs to taxonomy. Input taxonomy files must be TAB-delimited files where rows are either in the form:

1. SEQID[TAB]k\_\_Bacteria;p\_\_Firmicutes;c\_\_Clostridia;o\_\_Clostridiales;f\_\_;  
g\_\_;
2. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales;;;
3. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales
4. SEQID[TAB]D\_0\_\_Bacteria;D\_1\_\_Firmicutes;D\_2\_\_Clostridia;  
D\_3\_\_Clostridiales;D\_4\_\_;D\_5\_\_;

Compatible taxonomy files are in:

- Greengenes (<http://greengenes.secondgenome.com/downloads>);
- QIIME-formatted SILVA (<https://www.arb-silva.de/download/archive/qiime/>);
- UNITE (<https://unite.ut.ee/repository.php>);
- Human Oral Microbiome Database (HOMD) (<http://www.homd.org/>).

The output taxonomy file returned by `classify` is a TAB-delimited file where each row is always in the format:

```
SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales
```

## OTU table and taxonomy tables

The OTU table returned by *otu* is an OTU x sample, TAB-delimited text file, containing the number of times an OTU is found in each sample:

```
OTU      Mw_01 Mw_02 Mw_03 ...
DENOVO1 151   178   177   ...
DENOVO2 339   181   142   ...
DENOVO3 533   305   63    ...
DENOVO4 166   299   115   ...
...      ...   ...   ...   ...
```

The *tabletotax* command returns the “taxonomy tables” for each taxonomic level, e.g.:

```
OTU
Bacteria;Bacteroidetes      Mw_01 Mw_02 Mw_03 ...
1363 1543 1168 ...
Bacteria;Cyanobacteria/Chloroplast 0 0 0 ...
Bacteria;Firmicutes        6257 5780 6761 ...
Bacteria;Lentisphaerae     0 1 0 ...
...                          ...  ...  ...  ...
```

## Sample data

The sample data file contains all of the information about the samples. In QIIME this file is called [Mapping File](#). In micca, the sample data file must be a TAB-delimited text file (a row for each sample). The first column must be the sample identifier (assigned in *merge*, *split* or *mergepairs*):

```
ID      Group Altitude
Mw_01   Mw1    492
Mw_02   Mw1    492
Mw_09   Mw1    492
Mw_12   Mw1    492
...     ...    ...
```

## Phylogenetic tree

Only the [Newick format](#) is supported.



### Version 1.6.2 (bug fix release)

- Definitely fix the “new-line error” in classify
- Fix bar plots in “stats”

### Version 1.6.1 (bug fix release)

- Fix the “new-line error” in classify

### Version 1.6.0

- swarm updated to version 2.1.12;
- Now the mergepairs command allows merging staggered reads by default. With the new option `-n/--nostagger` the command produces the same results of the previous versions ( $\leq 1.5$ );
- classify and tabletotax commands now strip the `D_X__` prefix from the Silva taxonomy files;
- Documentation updated;
- Fix: remove duplicate file closing in `micca.api.merge()`.

### Version 1.5.0

- Now the NAST algorithm trims candidate sequences to that which is bound by the beginning and end points of the alignment span; with the new option `--nast-notrim` in `micca msa` produces the same results of the previous version ( $\leq 1.4.0$ );

- Y-scale in `micca tablebar` when `--raw` fixed;
- “text file busy error” in `micca msa (nast)` fixed;
- pandas deprecation warnings fixed;
- documentation updated.

## Version 1.4.0

- No-filter option added to `micca.api.msa.nast()` (do not remove positions which are gaps in every sequence) and to the `msa` command (`--nast-nofilter` option);
- documentation improved.

## Version 1.3.0

- Swarm clustering algorithm added to `micca otu`;
- `micca.api.otu.denovo_swarm()` function added;
- micca v1.3.0 includes: VSEARCH v1.9.5, MUSCLE v3.8.31, FastTree v2.1.8, swarm v2.1.8;
- documentation updated.

## Version 1.2.2

- Now micca can generate plots with matplotlib when the `DISPLAY` environment variable is undefined;
- `MANIFEST.in`, Dockerfile updated.

## Version 1.2.1

- Dockerfile added;
- Documentation improved.

## Version 1.2.0

- Hits output file option added to `micca.api.msa.nast()` (`hits_fn` option) and to the `msa` command (`--nast-hits` option);
- `setup.py` improved.

## Version 1.1.0

- Strand option added in `classify` (consensus-based classifier), `msa` (NAST) and `otu` (closed-reference and open reference OTU picking protocols) commands. Now these commands search both strand (default) instead of the plus strand only.

## Version 1.0.0

- micca 1.0.0 includes: VSEARCH v1.9.5, MUSCLE v3.8.31, FastTree v2.1.8.



```
usage: micca classify [-h] -i FILE -o FILE [-m {cons,rdp,otuid}] [-r FILE]
                        [-x FILE] [--cons-id CONS_ID]
                        [--cons-maxhits CONS_MAXHITS]
                        [--cons-minfrac CONS_MINFRAC]
                        [--cons-mincov CONS_MINCOV] [--cons-strand {both,plus}]
                        [--cons-threads THREADS]
                        [--rdp-gene {16srrna,funallsu,fungalits_warcup,fungalits_unite}]
                        [--rdp-maxmem GB] [--rdp-minconf RDP_MINCONF]
```

micca classify assigns taxonomy for each sequence in the input file and provides three methods for classification:

\* VSEARCH-based consensus classifier (cons): input sequences are searched in the reference database with VSEARCH (<https://github.com/torognes/vsearch>). For each query sequence the method retrieves up to 'cons-maxhits' hits (i.e. identity >= 'cons-id'). Then, the most specific taxonomic label that is associated with at least 'cons-minfrac' of the hits is assigned. The method is similar to the UCLUST-based consensus taxonomy assigner presented in doi: 10.7287/peerj.preprints.934v2 and available in QIIME.

\* RDP classifier (rdp): only RDP classifier version >= 2.8 is supported (doi:10.1128/AEM.00062-07). In order to use this classifier RDP must be installed (download at <http://sourceforge.net/projects/rdp-classifier/files/rdp-classifier/>) and the RDPPATH environment variable setted. The available databases (--rdp-gene) are:

- 16S (16srrna)
- Fungal LSU (28S) (funallsu)
- Warcup ITS (fungalits\_warcup, doi: 10.3852/14-293)
- UNITE ITS (fungalits\_unite)

For more information about the RDP classifier go to

```
http://rdp.cme.msu.edu/classifier/classifier.jsp
```

\* OTU ID classifier (otuid): simply perform a sequence ID matching with the reference taxonomy file. Recommended strategy when the closed reference clustering (`--method closedref` in `micca-otu`) was performed. OTU ID classifier requires a tab-delimited file where the first column contains the current OTU ids and the second column the reference taxonomy ids (see `otuids.txt` in `micca-otu`), e.g.:

```
REF1[TAB]1110191
REF2[TAB]1104777
REF3[TAB]1078527
...
```

The input reference taxonomy file (`--ref-tax`) should be a tab-delimited file where rows are either in the form:

1. SEQID[TAB]k\_\_Bacteria;p\_\_Firmicutes;c\_\_Clostridia;o\_\_Clostridiales;f\_\_;g\_\_;
2. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales;;;
3. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales
4. SEQID[TAB]D\_0\_\_Bacteria;D\_1\_\_Firmicutes;D\_2\_\_Clostridia;D\_3\_\_Clostridiales;D\_4\_\_;D\_5\_\_;

Compatible reference database are Greengenes (<http://greengenes.secondgenome.com/downloads>), QIIME-formatted SILVA (<https://www.arb-silva.de/download/archive/qiime/>) and UNITE (<https://unite.ut.ee/repository.php>).

The output file is a tab-delimited file where each row is in the format:

```
SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales
```

optional arguments:

```
-h, --help          show this help message and exit
```

arguments:

```
-i FILE, --input FILE          input FASTA file (for 'cons' and 'rdp') or a tab-delimited OTU ids file (for 'otuid') (required).
-o FILE, --output FILE         output taxonomy file (required).
-m {cons,rdp,otuid}, --method {cons,rdp,otuid}
                                classification method (default cons)
-r FILE, --ref FILE            reference sequences in FASTA format, required for 'cons' classifier.
-x FILE, --ref-tax FILE        tab-separated reference taxonomy file, required for 'cons' and 'otuid' classifiers.
```

VSEARCH-based consensus classifierspecific options:

```
--cons-id CONS_ID            sequence identity threshold (0.0 to 1.0, default 0.9).
--cons-maxhits CONS_MAXHITS  number of hits to consider (>=1, default 3).
--cons-minfrac CONS_MINFRAC  for each taxonomic rank, a specific taxa will be assigned if it is present in at least MINFRAC of the hits (0.0 to 1.0, default 0.5).
```

```
--cons-mincov CONS_MINCOV
    reject sequence if the fraction of alignment to the
    reference sequence is lower than MINCOV. This
    parameter prevents low-coverage alignments at the end
    of the sequences (default 0.75).
--cons-strand {both,plus}
    search both strands or the plus strand only (default
    both).
--cons-threads THREADS
    number of threads to use (1 to 256, default 1).

RDP Classifier/Database specific options:
--rdp-gene {16srrna, fungallsu, fungalits_warcup, fungalits_unite}
    marker gene/region
--rdp-maxmem GB
    maximum memory size for the java virtual machine in GB
    (default 2)
--rdp-minconf RDP_MINCONF
    minimum confidence value to assign taxonomy to a
    sequence (default 0.8)
```

#### Examples

Classification of 16S sequences using the consensus classifier and Greengenes:

```
micca classify -m cons -i input.fasta -o tax.txt \
--ref greengenes_2013_05/rep_set/97_otus.fasta \
--ref-tax greengenes_2013_05/taxonomy/97_otu_taxonomy.txt
```

Classification of ITS sequences using the RDP classifier and the UNITE database:

```
micca classify -m rdp --rdp-gene fungalits_unite -i input.fasta \
-o tax.txt
```

OTU ID matching after the closed reference OTU picking protocol:

```
micca classify -m otuid -i otuids.txt -o tax.txt \
--ref-tax greengenes_2013_05/taxonomy/97_otu_taxonomy.txt
```





```
usage: micca convert [-h] -i FILE -o FILE [-q FILE] [-d DEFAULTQ]
                  [-f INPUT_FORMAT] [-F OUTPUT_FORMAT]

micca convert converts between sequence file formats. See
http://biopython.org/wiki/SeqIO#File\_Formats for a comprehensive list
of the supported file formats.

Supported input formats:
abi, abi-trim, ace, embl, embl-cds, fasta, fasta-qual, fastq, fastq-illumina,
fastq-sanger, fastq-solexa, gb, genbank, genbank-cds, ig, imgt, pdb-atom,
pdb-seqres, phd, pir, qual, seqxml, sff, sff-trim, swiss, tab, uniprot-xml

Supported output formats:
embl, fasta, fastq, fastq-illumina, fastq-sanger, fastq-solexa, gb, genbank,
imgt, phd, qual, seqxml, sff, tab

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE          input sequence file (required).
-o FILE, --output FILE        output sequence file (required).
-q FILE, --qual FILE          input quality file (required for 'fasta-qual' input
                              format).
-d DEFAULTQ, --defaultq DEFAULTQ
                              default phred quality score for format-without-quality
                              to format-with-quality conversion (default 40).
-f INPUT_FORMAT, --input-format INPUT_FORMAT
                              input file format (default fastq).
-F OUTPUT_FORMAT, --output-format OUTPUT_FORMAT
                              input file format (default fasta).

Examples
```

Convert FASTA+QUAL files into a FASTQ (Sanger/Illumina 1.8+) file:

```
micca convert -i input.fasta -q input.qual -o output.fastq \  
-f fasta-qual -F fastq
```

Convert a SFF file into a FASTQ (Sanger/Illumina 1.8+) file:

```
micca convert -i input.sff -o output.fastq -f sff -F fastq
```

# CHAPTER 15

---

## filter

---

```
usage: micca filter [-h] -i FILE -o FILE [-e MAXEERATE] [-m MINLEN] [-t]
                    [-n MAXNS] [-f {fastq,fasta}]
```

micca filter filters sequences according to the maximum allowed expected error (EE) rate %. Optionally, you can:

- \* discard sequences that are shorter than the specified length (suggested for Illumina overlapping paired-end (already merged) reads) (option --minlen MINLEN);

- \* discard sequences that are shorter than the specified length AND truncate sequences that are longer (suggested for Illumina and 454 unpaired reads) (options --minlen MINLEN --trunc);

- \* discard sequences that contain more than a specified number of Ns (--maxns).

Sequences are first shortened and then filtered. Overlapping paired reads with should be merged first (using micca-mergepairs) and then filtered.

The expected error (EE) rate % in a sequence of length L is defined as (doi: 10.1093/bioinformatics/btv401):

$$\text{EE rate \%} = \frac{\text{sum(error probabilities)}}{L} * 100$$

Before filtering, run 'micca filterstats' to see how many reads will pass the filter at different minimum lengths with or without truncation, given a maximum allowed expected error rate % and maximum allowed number of Ns.

micca-filter is based on VSEARCH (<https://github.com/torognes/vsearch>).

```
optional arguments:
-h, --help                show this help message and exit

arguments:
-i FILE, --input FILE      input FASTQ file, Sanger/Illumina 1.8+ format
                           (phred+33) (required).
-o FILE, --output FILE     output FASTA/FASTQ file (required).
-e MAXEERATE, --maxeerate MAXEERATE
                           discard sequences with more than the specified expected
                           error rate % (values <=1%, i.e. less or equal than one
                           error per 100 bases, are highly recommended).
                           Sequences are discarded after truncation (if enabled)
                           (default 1).
-m MINLEN, --minlen MINLEN
                           discard sequences that are shorter than MINLEN
                           (default 1).
-t, --trunc               truncate sequences that are longer than MINLEN
                           (disabled by default).
-n MAXNS, --maxns MAXNS   discard sequences with more than the specified number
                           of Ns. Sequences are discarded after truncation
                           (disabled by default).
-f {fastq,fasta}, --output-format {fastq,fasta}
                           file format (default fasta).
```

#### Examples

Truncate reads at 300 bp, discard low quality sequences  
(with EE rate > 0.5%) and write a FASTA file:

```
micca filter -i reads.fastq -o filtered.fasta -m 300 -t -e 0.5
```

# CHAPTER 16

---

## filterstats

---

```
usage: micca filterstats [-h] -i FILE [-o DIR] [-t TOPN]
                        [-e MAXEERATES [MAXEERATES ...]] [-n MAXNS]

micca filterstats reports the fraction of reads that would pass for each
specified maximum expected error (EE) rate %% and the maximum number of
allowed Ns after:

* discarding sequences that are shorter than the specified length
(suggested for Illumina overlapping paired-end (already merged)
reads);

* discarding sequences that are shorter than the specified length AND
truncating sequences that are longer (suggested for Illumina and 454
unpaired reads);

Parameters for the 'micca filter' command should be chosen for each
sequencing run using this tool.

micca filterstats returns in the output directory 3 files:

* filterstats_minlen.txt: fraction of reads that would pass the filter after
the minimum length filtering;
* filterstats_trunclen.txt: fraction of reads that would pass the filter after
the minimum length filtering + truncation;
* filterstats_plot.png: plot in PNG format.

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE      input FASTQ file, Sanger/Illumina 1.8+ format
                           (phred+33) (required).
-o DIR, --output DIR     output directory (default .).
-t TOPN, --topn TOPN     perform statistics on the first TOPN sequences
```

```
                                (disabled by default)
-e MAXEERATES [MAXEERATES ...], --maxeeres MAXEERATES [MAXEERATES ...]
                                max expected error rates (%). (default [0.25, 0.5,
                                0.75, 1, 1.25, 1.5])
-n MAXNS, --maxns MAXNS
                                max number of Ns. (disabled by default).
```

#### Examples

Compute filter statistics on the top 10000 sequences, predicting the fraction of reads that would pass for each maximum EE error rate (default values):

```
micca filterstats -i input.fastq -o stats -t 10000
```

# CHAPTER 17

---

## merge

---

```
usage: micca merge [-h] -i FILE [FILE ...] -o FILE [-s SEP] [-f {fastq,fasta}]
```

micca merge merges several FASTQ or FASTA files in a single file. Different samples will be merged in a single file and sample names will be appended to the sequence identifier (e.g. >SEQID;sample=SAMPLENAME). Sample names are defined as the leftmost part of the file name splitted by the first occurrence of '.' (-s/--sep option). Whitespace characters in names will be replaced with a single character underscore ('\_').

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE [FILE ...], --input FILE [FILE ...]  
input FASTQ/FASTA file(s) (required).

-o FILE, --output FILE  
output FASTQ/FASTA file (required).

-s SEP, --sep SEP Sample names are defined as the leftmost part of the file name splitted by the first occurrence of 'SEP' (default .)

-f {fastq,fasta}, --format {fastq,fasta}  
file format (default fastq).

Examples

Merge files in FASTA format:

```
micca merge -i in1.fasta in2.fasta in3.fasta -o merged.fasta \  
-f fasta
```





```
usage: micca mergepairs [-h] -i FILE [FILE ...] -o FILE [-r FILE]
                        [-l MINOVLEN] [-d MAXDIFFS] [-p PATTERN] [-e REPL]
                        [-s SEP] [-n] [--notmerged-fwd FILE]
                        [--notmerged-rev FILE]
```

micca mergepairs merges paired-end sequence reads into one sequence.

A single merging of a pair of FASTQ files can be simply performed using both `-i/--input` and `-r/--reverse` options.

When the option `-r/--reverse` is not specified:

1. you can indicate several forward files (with the option `-i/--input`);

2. the reverse file name will be constructed by replacing the string `'_R1'` in the forward file name with `'_R2'` (typical in Illumina file names, see options `-p/--pattern` and `-e/--repl`);

3. after the merging of the paired reads, different samples will be merged in a single file and sample names will be appended to the sequence identifier (e.g. `>SEQID;sample=SAMPLENAME`), as in 'micca merge' and 'micca split'. Sample names are defined as the leftmost part of the file name splitted by the first occurrence of `'_'` (`-s/--sep` option). Whitespace characters in names will be replaced with a single character underscore (`'_'`).

micca mergepairs wraps VSEARCH (<https://github.com/torognes/vsearch>). Statistical testing of significance is performed in a way similar to PEAR (doi: 10.1093/bioinformatics/btt593). The quality of merged bases is computed as in USEARCH (doi: 10.1093/bioinformatics/btv401).

By default staggered read pairs (staggered pairs are pairs where the 3' end of the reverse read has an overhang to the left of the 5' end of the forward read) will be merged. To override this feature (and therefore to discard staggered alignments) set the `-n/--nostagger`

option.

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE [FILE ...], --input FILE [FILE ...]  
forward FASTQ file(s), Sanger/Illumina 1.8+ format  
(phred+33) (required).

-o FILE, --output FILE  
output FASTQ file (required).

-r FILE, --reverse FILE  
reverse FASTQ file, Sanger/Illumina 1.8+ format  
(phred+33).

-l MINOVLEN, --minovlen MINOVLEN  
minimum overlap length (default 32).

-d MAXDIFFS, --maxdiffs MAXDIFFS  
maximum number of allowed mismatches in the overlap  
region (default 8).

-p PATTERN, --pattern PATTERN  
when the reverse filename is not specified, it will be  
constructed by replacing 'PATTERN' in the forward file  
name with 'REPL' (default \_R1).

-e REPL, --repl REPL  
when the reverse filename is not specified, it will be  
constructed by replacing 'PATTERN' in the forward file  
name with 'REPL' (default \_R2).

-s SEP, --sep SEP  
when the reverse file name is not specified, sample  
names are appended to the sequence identifier (e.g.  
>SEQID;sample=SAMPLENAME). Sample names are defined as  
the leftmost part of the file name splitted by the  
first occurrence of 'SEP' (default \_)

-n, --nostagger  
forbid the merging of staggered read pairs. Without  
this option the command will merge staggered read  
pairs and the 3' overhang of the reverse read will be  
not included in the merged sequence.

--notmerged-fwd FILE write not merged forward reads.  
--notmerged-rev FILE write not merged reverse reads.

Examples

Merge reads with a minimum overlap length of 50 and maximum number  
of allowed mismatches of 3:

```
micca mergepairs -i reads1.fastq -r reads2.fastq -o merged.fastq \
-l 50 -d 3
```

Merge several illumina paired reads (typically named \*\_R1\*.fastq and  
\*\_R2\*.fastq):

```
micca mergepairs -i *_R1*.fastq -o merged.fastq --notmerged-fwd \
notmerged_fwd.fastq --notmerged-rev notmerged_rev.fastq
```

```
usage: micca msa [-h] -i FILE -o FILE [-m {muscle,nast}]
               [--muscle-maxiters MUSCLE_MAXITERS] [--nast-template FILE]
               [--nast-id NAST_ID] [--nast-threads NAST_THREADS]
               [--nast-mincov NAST_MINCOV] [--nast-strand {both,plus}]
               [--nast-notaligned FILE] [--nast-hits FILE] [--nast-nofilter]
               [--nast-notrim]
```

micca msa performs a multiple sequence alignment (MSA) on the input file in FASTA format. micca msa provides two approaches for MSA:

\* MUSCLE (doi: 10.1093/nar/gkh340). It is one of the most widely-used multiple sequence alignment software;

\* Nearest Alignment Space Termination (NAST) (doi: 10.1093/nar/gkl244). MICCA provides a very fast and memory efficient implementation of the NAST algorithm. The algorithm is based on VSEARCH (<https://github.com/torognes/vsearch>). It requires a pre-aligned database of sequences (--nast-template). For 16S data, a good template file is the Greengenes Core Set ([http://greengenes.lbl.gov/Download/Sequence\\_Data/Fasta\\_data\\_files/core\\_set\\_aligned.fasta.imputed](http://greengenes.lbl.gov/Download/Sequence_Data/Fasta_data_files/core_set_aligned.fasta.imputed)).

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE, --input FILE input FASTA file (required).

-o FILE, --output FILE output MSA file in FASTA format (required).

-m {muscle,nast}, --method {muscle,nast} multiple sequence alignment method (default muscle).

MUSCLE specific options:

--muscle-maxiters MUSCLE\_MAXITERS

maximum number of MUSCLE iterations. Set to 2 for a good compromise between speed and accuracy (>=1 default 16).

NAST specific options:

```
--nast-template FILE  multiple sequence alignment template file in FASTA
                       format.
--nast-id NAST_ID     sequence identity threshold to consider a sequence a
                       match (0.0 to 1.0, default 0.75).
--nast-threads NAST_THREADS
                       number of threads to use (1 to 256, default 1).
--nast-mincov NAST_MINCOV
                       reject sequence if the fraction of alignment to the
                       template sequence is lower than MINCOV. This parameter
                       prevents low-coverage alignments at the end of the
                       sequences (default 0.75).
--nast-strand {both,plus}
                       search both strands or the plus strand only (default
                       both).
--nast-notaligned FILE
                       write not aligned sequences in FASTA format.
--nast-hits FILE      write hits on a TAB delimited file with the query
                       sequence id, the template sequence id and the
                       identity.
--nast-nofilter       do not remove positions which are gaps in every
                       sequenceces (useful if you want to apply a Lane mask
                       filter before the tree inference).
--nast-notrim         force to align the entire candidate sequence (i.e. do
                       not trim the candidate sequence to that which is bound
                       by the beginning and end points of of the alignment
                       span
```

Examples

De novo MSA using MUSCLE:

```
micca msa -i input.fasta -o msa.fasta
```

Template-based MSA using NAST, the Greengenes alignment as template (clustered at 97% similarity) 4 threads and a sequence identity threshold of 75%:

```
micca msa -i input.fasta -o msa.fasta -m nast --nast-threads 4 \
--nast-template greengenes_2013_05/rep_set_aligned/97_otus.fasta
```

```
usage: micca otu [-h] -i FILE [-o DIR] [-r FILE]
                [-m {denovo_greedy,denovo_swarm,closed_ref,open_ref}] [-d ID]
                [-n MINCOV] [-t THREADS] [-c] [-g {dgc,agc}] [-s MINSIZE]
                [-a {both,plus}] [--swarm-differences SWARM_DIFFERENCES]
                [--swarm-fastidious]
```

micca otu assigns similar sequences (marker genes such as 16S rRNA and the fungal ITS region) to operational taxonomic units (OTUs).

Trimming the sequences to a fixed position before clustering is *strongly recommended* when they cover partial amplicons or if quality deteriorates towards the end (common when you have long amplicons and single-end sequencing).

Removing ambiguous nucleotides 'N' (with the option `--maxns 0` in micca filter) is mandatory if you use the de novo swarm clustering method.

micca otu provides the following protocols:

\* de novo greedy clustering (denovo\_greedy): sequences are clustered without relying on an external reference database, using an approach similar to the UPARSE pipeline (doi: 10.1038/nmeth.2604): i) predict sequence abundances of each sequence by dereplication; ii) greedy clustering; iii) remove chimeric sequences (de novo, optional) from the representatives; iv) map sequences to the representatives.

\* de novo swarm (denovo\_swarm): sequences are clustered without relying on an external reference database, using swarm (doi: 10.7717/peerj.593, doi: 10.7717/peerj.1420, <https://github.com/torognes/swarm>): i) predict sequence abundances of each sequence by dereplication; ii) swarm clustering; iii) remove chimeric sequences (de novo, optional) from the representatives.

\* closed-reference clustering (closed\_ref): sequences are clustered against an external reference database and reads that could not be

matched are discarded.

\* open-reference clustering (open\_ref): sequences are clustered against an external reference database and reads that could not be matched are clustered with the 'de novo greedy' protocol.

Outputs:

\* otutable.txt: OTU x sample, TAB-separated OTU table file, containing the number of times an OTU is found in each sample.

\* otus.fasta: FASTA file containing the representative sequences (OTUs);

\* otuids.txt: OTU ids to original sequence ids (tab-delimited text file)

\* hits.txt: three-columns, TAB-separated file:

1. matching sequence
2. representative (seed)
3. identity (if available, else '\*'), defined as:

```

        matching columns
        ----- ;
        alignment length - terminal gaps

```

\* otuschim.fasta (only for 'denovo\_greedy', 'denovo\_swarm' and 'open\_ref' when --rmchim is specified): FASTA file containing the chimeric otus;

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE, --input FILE input fasta file (required).

-o DIR, --output DIR output directory (default .).

-r FILE, --ref FILE reference sequences in fasta format, required for 'closed\_ref' and 'open\_ref' clustering methods.

-m {denovo\_greedy,denovo\_swarm,closed\_ref,open\_ref}, --method {denovo\_greedy,denovo\_swarm,closed\_ref,open\_ref} clustering method (default denovo\_greedy)

-d ID, --id ID sequence identity threshold (for 'denovo\_greedy', 'closed\_ref' and 'open\_ref', 0.0 to 1.0, default 0.97).

-n MINCOV, --mincov MINCOV reject sequence if the fraction of alignment to the reference sequence is lower than MINCOV. This parameter prevents low-coverage alignments at the end of the sequences (for 'closed\_ref' and 'open\_ref' clustering methods, default 0.75).

-t THREADS, --threads THREADS number of threads to use (1 to 256, default 1).

-c, --rmchim remove chimeric sequences (for 'denovo\_greedy', 'denovo\_swarm' and 'open\_ref' OTU picking methods).

-g {dgc,agc}, --greedy {dgc,agc} greedy clustering strategy, distance (DGC) or abundance-based (AGC) (for 'denovo\_greedy' and 'open\_ref' clustering methods) (default dgc).

```
-s MINSIZE, --minsize MINSIZE
    discard sequences with an abundance value smaller than
    MINSIZE after dereplication (>=1, default 2).
    Recommended value is 2 (i.e. discard singletons) for
    'denovo_greedy' and 'open_ref', 1 (do not discard
    anything) for 'denovo_swarm'.
-a {both,plus}, --strand {both,plus}
    search both strands or the plus strand only (for
    'closed_ref' and 'open_ref' clustering methods,
    default both).
```

#### Swarm specific options:

```
--swarm-differences SWARM_DIFFERENCES
    maximum number of differences allowed between two
    amplicons. Commonly used d values are 1 (linear
    complexity algorithm), 2 or 3, rarely higher. (>=0,
    default 1).
--swarm-fastidious
    when working with SWARM_DIFFERENCES=1, perform a
    second clustering pass to reduce the number of small
    OTUs (recommended option).
```

#### Examples

De novo clustering with a 97% similarity threshold and remove chimeric OTUs:

```
micca otu -i input.fasta --method denovo_greedy --id 0.97 -c
```

Open-reference OTU picking protocol with a 97% similarity threshold, without removing chimeras in the de novo protocol step and using 8 threads:

```
micca otu -i input.fasta --method open_ref --threads 8 --id 0.97 \
--ref greengenes_2013_05/rep_set/97_otus.fasta
```

De novo swarm clustering with the protocol suggested by the authors using 4 threads (see <https://github.com/torognes/swarm> and <https://github.com/torognes/swarm/wiki>):

```
micca otu -i input.fasta --method denovo_swarm --threads 4 \
--swarm-fastidious --rmchim --minsize 1
```





```
usage: micca root [-h] -i FILE -o FILE [-m {midpoint,outgroup}]
               [-t TARGETS [TARGETS ...]]

micca root reroot the input tree:

* at the calculated midpoint between the two most distant tips of the
tree (--method midpoint);

* with the outgroup clade containing the given taxa (leaf nodes),
i.e. the common ancestor of the outgroup (--method outgroup).

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE      input FASTA file (required).
-o FILE, --output FILE    output MSA file in FASTA format (required).
-m {midpoint,outgroup}, --method {midpoint,outgroup}
                           rooting method (default midpoint).
-t TARGETS [TARGETS ...], --targets TARGETS [TARGETS ...]
                           list of targets defining the outgroup (required for
                           the outgroup method).
```

#### Examples

##### Midpoint rooting:

```
micca root -i input.tree -o input_rooted.tree
```

##### Rooting with outgroup:

```
micca root -i input.tree -o input_rooted.tree -m outgroup DENOVO1 DENOVO2
```



```
usage: micca split [-h] -i FILE -o FILE -b FILE [-n FILE] [-c FILE] [-s N]
                [-e MAXE] [-t] [-f {fastq,fasta}]
```

micca split assign the multiplexed reads to samples based on their 5' nucleotide barcode (demultiplexing) provided by the FASTA file (`--barcode`). micca split creates a single FASTQ or FASTA file with sample information (e.g. `>SEQID;sample=SAMPLENAME`) appended to the sequence identifier. Barcode and the sequence preceding it is removed by default, e.g.:

```
Barcode file:      Input file:
>SAMPLE1           >SEQ1
TCAGTCAG           TCAGTCAGGCCACGGCTAACTAC...
...                ...
```

the output will be:

```
>SEQ1;sample=SAMPLE1
GCCACGGCTAACTAC...
...
```

optional arguments:

```
-h, --help          show this help message and exit
```

arguments:

```
-i FILE, --input FILE          input FASTQ/FASTA file (required).
-o FILE, --output FILE         output FASTQ/FASTA file (required).
-b FILE, --barcode FILE        barcode file in FASTA format (required).
-n FILE, --notmatched FILE     write reads in which no barcode was found.
-c FILE, --counts FILE
```

```
write barcode counts in a tab-delimited file.
-s N, --skip N      skip N bases before barcode matching (e.g. if your
                    sequences start with the control sequence 'TCAG'
                    followed by the barcode, set to 4) (>=0, default 0).
-e MAXE, --maxe MAXE maximum number of allowed errors (>=0, default 1).
-t, --notrim       do not trim barcodes and the sequence preceding it
                    from sequences.
-f {fastq,fasta}, --format {fastq,fasta}
                    file format (default fastq).
```

#### Examples

Split 'reads.fastq' and write the notmatched sequences in the file 'notmatched.fastq':

```
micca split -i input.fastq -o splitted.fastq -b barcode.fasta \
-n notmatched.fastq
```

```
usage: micca stats [-h] -i FILE [-o DIR] [-n TOPN]

micca stats reports statistics on reads in a FASTQ file. micca stats
returns in the output directory 3 tab-delimited text files:

* stats_lendist.txt: length distribution;
* stats_qualdist.txt: Q score distribution;
* stats_qualsumm.txt: quality summary. For each read position, the
following statistics are reported:
- L: read position;
- NPctCum: percent of reads with at least L bases;
- QAv: average Q score at position L;
- EERatePctAv: average expected error (EE) rate %.

Moreover, micca stats returns the respective plots in PNG format,
stats_lendist_plot.png, stats_qualdist_plot.png, and
stats_qualsumm_plot.png.

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE          input FASTQ file, Sanger/Illumina 1.8+ format
                                (phred+33) (required).
-o DIR, --output DIR         output directory (default .).
-n TOPN, --topn TOPN         perform statistics only on the first TOPN sequences
                                (disabled by default).

Examples

Compute statistics on the top 10000 sequences of input.fastq:

    micca stats -i input.fastq -o stats -n 10000
```



```
usage: micca tablebar [-h] -i FILE -o FILE [-r] [-t N] [--xticklabelsize SIZE]
                        [-f {pgf,ps,svg,rgba,raw,svgz,pdf,eps,png}]

micca tablebar generates a relative abundance bar plot from
OTU or taxa tables. The table must be an OTU/taxon x sample,
TAB-separated file (see 'micca otu').

optional arguments:
-h, --help                show this help message and exit

arguments:
-i FILE, --input FILE      input OTU table file (required).
-o FILE, --output FILE     output image file (required).
-r, --raw                  plot raw values (i.e. counts) instead of relative
                           abundances.
-t N, --topn N             plot the top N abundant taxa (default 12).
--xticklabelsize SIZE     x tick label size (default 8).
-f {pgf,ps,svg,rgba,raw,svgz,pdf,eps,png}, --format {pgf,ps,svg,rgba,raw,svgz,pdf,eps,
↪png}                      output file format (default png).

Example

micca tablebar -i otutable.txt -o otutable_plot.png
```





---

## tablerare

---

```
usage: micca tablerare [-h] -i FILE -o FILE -d DEPTH [-r] [-s SEED]
```

Rarefy an OTU table by subsampling, with or without replacement. Samples that have fewer counts than the depth are omitted from the output table. OTUs that are not present in at least one sample are omitted from the output table.

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE, --input FILE input OTU table file (required).  
-o FILE, --output FILE output rarefied OTU table file (required).  
-d DEPTH, --depth DEPTH sample depth (>0, required).  
-r, --replace subsample with replacement.  
-s SEED, --seed SEED random seed (default 0).

Examples

Rarefy an OTU table at a depth of 1000 sequences/sample:

```
micca tablerare -i otutable.txt -o otutable_rare.txt -d 1000
```



---

## tablestats

---

```
usage: micca tablestats [-h] -i FILE [-o DIR] [-t STEP] [-r] [-s SEED]

micca tablestats reports a sample summary, an OTU summary and
the rarefaction curves for the input OTU table. The
rarefaction curves are evaluated using the interval of 'step'
(-t/--step) sample depths, always including 1 and the total
sample size.

micca filterstats returns in the output directory 4 files:

* tablestats_samplesumm.txt: samples summary;
* tablestats_otusumm.txt: OTUs summary;
* tablestats_rarecurve.txt: rarefaction curves in text format.
* tablestats_rarecurve_plot.txt: rarefaction curves in png format.

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE          input FASTQ file, Sanger/Illumina 1.8+ format
                                (phred+33) (required).
-o DIR, --output DIR         output directory (default .).
-t STEP, --step STEP        sample depth interval (for rarefaction curves, default
                                500).
-r, --replace                subsample with replacement (for rarefaction curves).
-s SEED, --seed SEED        random seed (for rarefaction curves, default 0).

Examples

Compute OTU table statistics on otutable.txt:

    micca tablestats -i otutable.txt -o tablestats
```



## tabletotax

```
usage: micca tabletotax [-h] -i FILE -t FILE [-o DIR]
```

Given an OTU table and a taxonomy file, micca tabletotax creates in the output directory a table for each taxonomic level (taxtable1.txt, ..., taxtableN.txt). OTU counts are summed together if they have the same taxonomy at the considered level.

The OTU table must be an OTU x sample, TAB-separated OTU table file (see 'micca otu'). The taxonomy file must be a tab-delimited file where rows are either in the form (see 'micca classify'):

1. SEQID[TAB]k\_\_Bacteria;p\_\_Firmicutes;c\_\_Clostridia;o\_\_Clostridiales;f\_\_;g\_\_;
2. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales;;;;
3. SEQID[TAB]Bacteria;Firmicutes;Clostridia;Clostridiales
4. SEQID[TAB]D\_0\_\_Bacteria;D\_1\_\_Firmicutes;D\_2\_\_Clostridia;D\_3\_\_Clostridiales;D\_4\_\_;D\_5\_\_;

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE, --input FILE input OTU table file (required).  
-t FILE, --tax FILE input taxonomy file (required).  
-o DIR, --output DIR output directory (default .).

Examples

```
micca tabletotax -i otutable.txt -t tax.txt -o taxtables
```



```
usage: micca tobiom [-h] -i FILE -o FILE [-t FILE] [-s FILE] [-u FILE]

micca tobiom converts the micca OTU table into BIOM Version
1.0 (JSON) format. Optionally, taxonomy and/or sample
information can be added. When you convert on
(closed-reference) OTU table for PICRUST, replace OTU IDs with
the original sequence IDs use the option -u/--otuids.

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE      input OTU table file (required).
-o FILE, --output FILE     output BIOM file Version 1.0 (JSON) (required).
-t FILE, --tax FILE       add taxonomy information from a taxonomy file.
-s FILE, --sampledata FILE add sample information from a sample data file.
-u FILE, --otuids FILE    replace OTU IDs with the original sequence IDs. Useful
                           when the closed-reference OTU picking protocol was
                           performed for PICRUST

Example

micca tobiom -i otutable.txt -o output.biom -t tax.txt -s sampledata.txt
```





```
usage: micca tree [-h] -i FILE -o FILE [-m {fasttree,muscle}] [--fasttree-gtr]
                [--fasttree-fastest]
                [--muscle-cluster {upgmb,upgma,neighborjoining}]
```

micca tree infers phylogenetic trees from alignments. It provides two methods:

- \* FastTree (doi: 10.1371/journal.pone.0009490);
- \* MUSCLE (doi: 10.1093/nar/gkl244).

optional arguments:

-h, --help show this help message and exit

arguments:

-i FILE, --input FILE input FASTA file (required).  
-o FILE, --output FILE output tree in Newick format (required).  
-m {fasttree,muscle}, --method {fasttree,muscle} tree inference method (default fasttree).

FastTree specific options:

--fasttree-gtr use the generalized time-reversible (GTR)+CAT model instead of Jukes-Cantor+CAT (default False).  
--fasttree-fastest speed up the neighbor joining phase and reduce memory usage recommended for >50,000 sequences) (default False).

MUSCLE specific options:

--muscle-cluster {upgmb,upgma,neighborjoining} clustering algorithm (default upgmb).

Examples

Tree inference using FastTree and the generalized time-reversible

(GTR) +CAT model:

```
micca tree -i input.fasta -o tree.tree --fasttree-gtr
```

```
usage: micca trim [-h] -i FILE -o FILE [-w FORWARD [FORWARD ...]]
                [-r REVERSE [REVERSE ...]] [-e MAXERATE] [-c] [-W] [-R]
                [-f {fastq,fasta}]

micca trim trims forward and reverse primers from a FASTQ/FASTA file
using Cutadapt (doi: 10.14806/ej.17.1.200) internally. Primer and the
sequence preceding (for forward) or succeeding (for reverse) it are
removed. Optionally, reads that do not contain the primers (untrimmed
reads) can be discarded with the options -W/--duforward and
-R/--dureverse. Recommended options are:

* always discard reads that do not contain the forward primer
(-W/--duforward option);

* for overlapping paired-end (already merged) reads, also discard
reads that do not contain the reverse primer (using both
-W/--duforward and -R/--dureverse options).

IUPAC codes and multiple primers are supported.

optional arguments:
-h, --help            show this help message and exit

arguments:
-i FILE, --input FILE      input FASTQ or FASTA file (required).
-o FILE, --output FILE    output FASTQ or FASTA file (required).
-w FORWARD [FORWARD ...], --forward FORWARD [FORWARD ...]
                          trim forward primer(s). Only the best matching primer
                          is removed.
-r REVERSE [REVERSE ...], --reverse REVERSE [REVERSE ...]
                          trim reverse primer(s). Only the best matching primer
                          is removed.
-e MAXERATE, --maxerate MAXERATE
```

```
                                maximum allowed error rate (default 0.1).
-c, --searchrc                  search reverse complement primers too (default False).
-W, --duforward                 discard untrimmed reads (reads that do not contain the
                                forward primer) (always recommended) (default False).
-R, --dureverse                 discard untrimmed reads (reads that do not contain the
                                reverse primer) (suggested option for overlapping
                                paired-end already merged reads) (default False).
-f {fastq,fasta}, --format {fastq,fasta}
                                file format (default fastq).
```

#### Examples

454 or Illumina single-end reads: trim forward primer and discard reads that do not contain it. Moreover, trim reverse primer:

```
micca trim -i input.fastq -o trimmed.fastq -w AGGATTAGATACCCTGGTA \  
-r CRRACGAGCTGACGAC -W
```

Illumina overlapping paired-end (already merged) reads: trim forward and reverse primers. Reads that do not contain the forward or the reverse primer will be discarded:

```
micca trim -i reads.fastq -o trimmed.fastq -w AGGATTAGATACCCTGGTA \  
-r CRRACGAGCTGACGAC -W -R
```